

FIG. 1

Constitutively Active Receptors

File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS A					
GROUP I					
MSHR_mouse	melanocyte-stimulating hormone	TMII	92 VSVIVGTTIIL K	adenylate cyclase activity/ HEK293, stably transfected	(Robbins, Nadeau et al. 1993)
	MSH				
CLASS A					
GROUP II					
5H1B_human	5-hydroxytryptamine _{1B}	C-terminus of IC3	313 REERKATKTLGI K, R, Q	binding of [³⁵ S]GTP[S]/ CHO-K1	(Pauwels, Gouble et al. 1999)
5H2A_human	5-hydroxytryptamine _{1A}	C-terminus of IC3	322 NEQRAGKVLGI K	IP production / COS-7	(Egan, Herrick-Davis et al. 1998)
2H2C_rat	5-hydroxytryptamine _{2C}	C-terminus of IC3	312 NEEDAGKVLGI L	PI hydrolysis / COS-7	(Herrick-Davis, Egan et al. 1997)

2/27

FIG. 1 (2 of 15)

FIG. 1 (2 of 15)

CLASS A GROUP II					
A1AB_human	α_{1B} -adrenergic alpha 1B-AR	TMDI	63 FAIVGQILVTL A	IP / COS-7	(Scheer, Fanelli et al. 1997)
A1AB_human	α_{1B} -adrenergic alpha 1B-AR	junction between TMDIII and IC2	142 CAISIDRYIGV A	IP / COS-7	(Scheer, Costa et al. 2000)
A1AB_human	α_{1B} -adrenergic alpha 1B-AR	junction between TMDIII and IC2	143 CAISIDRYIGV K	IP / COS-7	(Perez, Hwa et al. 1996)
A1AB_human	α_{1B} -adrenergic	TMDIII	128 AVDVLGCTASI F	IP / COS-1	
		carboxyl end of IC3	293 REKKAATLGI E	IP arachidonic acid release	(Hwa, Gaivin et al. 1997)
		TMV	204 EEFFYALFSSLG V	IP / COS-1	
A1AB_human	α_{1B} -adrenergic	C-terminal IC3	293 SREKKAAT X=19 different substitutions	PI / COS-7	(Kjelsberg, Cotechia et al. 1992)
A1AB_human	α_{1B} -adrenergic	C-terminus IC3	288 293 XFSREKKAATLGI K H L	PI hydrolysis / rat fibroblast	(Allen, Lefkowitz et al. 1991)
A2AA_human	α_{2C10} -adrenergic	C-terminal IC3 loop	373 (348?) EKRTFVLAV X=F, A, C, E, K	adenyl cyclase inhibition / HEK293	(Ren, Kurose et al. 1993)
ACM1_human	alpha-2AAR muscarinic Hm1	C-terminal IC3 loop junction	360 SVVGEKKAATLS A	PI / HEK(U293)	(Högger, Shockley et al. 1995)
ACM2-human	muscarinic acetylcholine M1 muscarinic acetylcholine M2	junction of IC3 and TMVI	390 KKVTRTLIA 1-4 A inserted	IP production, inhibition of cAMP production / COS-7	(Liu, Blin et al. 1996)

FIG. 1 (3 of 15)

CLASS A GROUP II ACM3_rat	m3 muscarinic (rat)	TMV1	507 TWTPYNYLVNT S	IP / COS-7	(Blüml, Mutschler et al. 1994)
ACM5_human	muscarinic acetylcholine M3	N-terminus to TMII	chimeric composed of m2 1-69 m5 77-445 m2 391-466	β -gal / NIH 3T3	(Burstein, Spalding et al. 1996)
ACM5_human	m5 muscarinic	TMV1	451 459 465 ALLA ELITW TPYNI MYLST N L H C V S F T	β -gal; radioligand binding / NIH-3T3	(Spalding, Burstein et al. 1998)
ACM5_human	muscarinic acetylcholine M5	junction of TMV1 and EC3	465 YNIWLVSTPCDKCV X=V,F,R,K,*more	β -gal; radioligand binding / NIH-3T3	(Spalding, Burstein et al. 1997)
BIAR_human	β -adrenergic	C-terminus	389 RKAFGLLCCA R	adenylyl cyclase; agonist binding / CHW	(Mason, Moore et al. 1999)
BZAR_human	β -adrenergic	C-terminal IC3 loop	266 272 FCLKFEKALKTGLI SR K A	adenylyl cyclase activation; agonist binding affinity / COS-7 or CHO	(Samama, Cotecchia et al. 1993); (Leftkowitz, Cotecchia et al. 1993)
DADR_human	dopamine D1A	carboxyl terminal IC3	264 SFPMSEFQETKVLKT I K 288 from D1B receptor	adenylyl cyclase; cAMP accumulation / HEK293	(Charpentier, Jarvie et al. 1996)
DADR_human	dopamine D1	TMV1	286 FVCCNLFFFIL A	CAMP accumulation / COS-7	(Cho, Taylor et al. 1996)
HH2R_rat	histamine H ₂	IC2	115 FMISLDRYCAV N.A.	cAMP production / HEK-293	(Alewijns, Timmerman et al. 2000)

FIG. 1 (4 of 15)

File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS A GROUP III OPSD_human	opsin				
	rhodopsin	TMII	⁹⁰ FWVLGGTSTLY D	transducin; phosphorylation by rhodopsin kinase / COS	(Rim and Oprea 1995)
		TMIII	113 GCVLGGFFAT Q		
		TMVII	292 296 MTIPAFFAKSAATY E G, E, N		
			²⁹² Ala neutral a.a converted to carboxylate and competes with ¹¹³ Glu for salt bridge with ²⁹⁶ Lys		
OPSD_human	opsin	TMIII	134 VVLATERYVVV I, Q, S	transducin; radioligand binding / COS	(Acharya and Kamik 1996)
OPSD_human	rhodopsin	TM6	257 RWVLTWIAFL Y, N	transducin, GTPγS uptake / COS	(Han, Smith et al. 1998)
OPSD_human	opsin	plus TM3 TMVII	plus GL13Q 296 PAFFAKSAATY G	transducin; radioligand binding / COS	(Govardhan and Oprea 1994); (Cohen, Yang et al. 1993)
	rhodopsin	IC2	X=E,M natural mutants + 10 different a.a. substitutions disrupts critical salt bridge between ²⁹⁶ Lys(TMVI) and ¹¹³ Glu(TMIII)		
			134 VVLATERYVVV Q		(Cohen, Yang et al. 1993)

FIG. 1 (6 of 15)

6/27

File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS A GROUP IV					
BRB2_human	bradykinin B ₂	TMIII	113 ATISQMLYSSI	IP production / COS-7	(Marie, Koch et al. 1999)
	B2 bradykinin BK-2	TMVI	A 256 LLFTICGLPFOI		
			F		

FIG. 1 (7 of 15)

File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS A GROUP V					
AG2R_rat	A _{1A} Type-1A angiotensin II	TMIII	111 ASVSFNFYASV A disrupts ¹¹¹ Aan(TMIII) - ¹¹¹ Tyr(TMIII) interaction	phospholipase C; IP production / COS-7	(Grobowski, Maigret et al. 1997)
AG2R_rat	A _{1A}	C-terminus of TM7 other multiple mutations	305 LFVGF ₃₀₅ AKKKF Q	IP production / HEK- 293; intracellular Ca ²⁺ mobilization / CHO	(Parnot, Bardin et al. 2000)
FM1R_human	Type-1A angiotensin II formylmethionylleucylphenylal- anine (fMLPR)	IC1	51 LVVWVFQFMHFTFTTISYLNKAVA LVVWVFQFMHFTFTTISYLNKAVA (K above conflicts with SWISS-PROT database)	PI production; phospholipase C stimulation / COS-7	(Amatunga, Dragas- Graonic et al. 1995)
IL8R_human	interleukin-8 receptor B CXCR-2 chemokine	IC2	138 ACISVDRYLAIVH V	IP production; Ca ²⁺ mobilization and actin polymerization / NIH 3T3	(Burger, Burger et al. 1999)
LSHR_human	luteinizing hormone (LH)	IC3	564 MATNDQTKIAKK G	cAMP production / HEK293	(Kudo, Otsuga et al. 1996)
LSHR_human	luteinizing hormone (LH)	TMVI	578 ILIFDTFCA G	cAMP production / COS-7	(Shenker, Laue et al. 1993)
LSHR_human	luteinizing hormone (LH)	TM6	571 577 KIARQKALLFTFTCH I I	cAMP production / COS-7	(Kosugi, Van Dop et al. 1995)
LSHR_rat	luteinizing hormone / human chorionic gonadotropin (LH/hCG)	TMVI	556 ILIFDTFCA G, Y	cAMP production / HEK 293T	(Bradbury, Kawate et al. 1997; Bradbury and Menon 1999)
OPRD_mouse	delta opioid receptor	TM3	128 KVLSDIYNNF A, K, H	adenylyl cyclase inhibition / COS-7	(Cavalli, Babey et al. 1999)
OXYR_human	oxytocin	IC2	137 LMSIDGCLATC A	IP production / COS-7	(Faneli, Barbier et al. 1999)

FIG. 1 (8 of 15)

PAFR_human	platelet-activating factor (PAF)	C-terminus of IC3	231 EYKRALMMVCTVLAV R	IP production / COS-7	Parent, Le Gouill et al. 1996
PAFR_human	platelet-activating factor (PAF)	TMIII	100 CLFFNYTCVS A	arachidonate release, IP production, adenylyl cyclase inhibition / CHO	(Ishii, Izumi et al. 1997)
PE23_human	prostaglandin E ₂ , EP3III EP3IV	C-terminal tail	360 FCOREFNG FCOMKRRRGRGSEFNG ↑truncated	inhibition of adenylyl cyclase / CHO-K1	(Jin, Mao et al. 1997)
PE23_mouse	prostaglandin E ₂ , EP3	carboxyl-terminal tail	336 KILLKKCOLRHT (3α) MRRHL (3β) ↑truncated	inhibition of adenylyl cyclase / CHO, stably expressed	(Hasegawa, Negishi et al. 1996)
THRR_human	thrombin	EC2 loop	259 268 CHDVNRETLLEGVYY DLKD KPF I 486	⁴⁵ Ca ²⁺ efflux, PI hydrolysis, reporter gene induction / COS-7	(Nanevicius, Wang et al. 1996)
TSHR_human	thyrotropin (TSHR) thyroid stimulating hormone	EC1	YTHALDQQTG F, M	inositol phosphate-- diacylglycerol cascade / COS-7	(Parma, Van Sande et al. 1995)
		EC2	568 YAKVSLCLPMD T		
TSHR_human	thyrotropin (TSHR) thyroid stimulating hormone	TMIII	509 ASELSVYTLTV A	adenylyl cyclase activation / COS-7	(Duprez, Parma et al. 1994)
		TMVII	672 YPLNSCANPEFL Y		
TSHR_human	thyrotropin (TSHR)	TMV	597 VAFVILCCCHV L	cAMP formation / COS-7 cells	(Esapa, Duprez et al. 1999)
TSHR_human	thyroid stimulating hormone thyrotropin (TSHR)	TMVII	677 CANPEFLAIFT V	cAMP formation / CHO cells	(Rusco, Wong et al. 1999)
TSHR_human	thyroid stimulating hormone thyrotropin (TSHR)	IC3	613 621 VRNPFYNSGDEDTIAK deletion	cAMP formation / COS-7	(Wonerow, Schonberg et al. 1998)

9/27

FIG. 1 (9 of 15)

TSHR_human	thyrotropin (TSHR)	IC3 / TMVI	623 632 KOTYAKEMAVLIETDFCN V I	cAMP activation / COS-7	(Paschke, Tonacchera et al. 1994)
V2R_human	thyroid stimulating hormone vasopressin V2	IC2	136 LAMTLQRHRAI A	cAMP formation / COS-7	(Morin, Cotte et al. 1998)

10/27

FIG. 1 (10 of 15)

File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS B GROUP I CALR_human	human calcitonin hCTR-1 hCTR-2	wild type (native) protein		adenylyl cyclase cAMP production / COS-1	(Cohen, Thaw et al. 1997)
CLASS B GROUP II PTRR_human	parathyroid hormone PTH/PTH-related peptide	junction of IC1 and TMII	223 TNTYIHWLFL R, K	cAMP accumulation / COS-7	(Schipani, Jensen et al. 1997)
		junction of IC3 and TMVI	410 KLKSTLVLP C, others		
CLASS B GROUP III GIPR_human	glucose-dependent insulinotropic peptide (GIP-R)	TMVI	340 VFAPVTEQAR P	cAMP production / L293	(Tseng and Lin 1997)
GLR_rat	glucagon	junction of IC loop I and TMII	178 TNTYIHWLFL R	cAMP accumulation / COS-7	(Hjorth, Orskov et al. 1998)
		IC end of TMVI	352 RLARSTLLIP A		
VIPR_human	vasoactive intestinal peptide 1 (VIP)	junction of IC loop I and TMII	178 RNTYIHWLFL R requires functional integrity of the N-terminal 8C domain	cAMP production / COS-7 or CHO	(Gaudin, Maoret et al. 1998) (Gaudin, Royer-Pessard et al. 1998)
		junction of IC loop 3 and TMVI	343 LARETLLIP X= R, P		

FIG. 1 (12 of 15)

File Name	Receptor	Mutation Site	Sequence	Assay / Cells	Reference
CLASS D O74283 RCB2 C. cinereus	pheromone	TM6	229 PLSAYQYHGT P	heterologous yeast assay	(Olesnicki, Brown et al. 1999)
STE2_yeast	pheromone α -factor	TM6	258 QSLIVESLIIFI LL	<i>lacZ</i> reporter gene	(Konopka, Margalit et al. 1996)
STE2_yeast	pheromone α -factor	double mutations TM5 and TM6	223 MSFVLVIVLILAIR C C 247 251 DSFHLINSCQSL CC CC	<i>lacZ</i> reporter gene / yeast	(Dube, DeCostanzo et al. 2000)
STE3_yeast	pheromone α -factor	IC3	double mutations shaded double mutations 194 DVRDLHCTNS Q	β -galactosidase	(Boone, Davis et al. 1993)
STE2_yeast	pheromone α -factor	TM6	253 258 LINSQSLIVESLIIFI L LP	β -galactosidase	(Somers, Martin et al. 2000)

FIG. 1 (13 of 15)

Bibliography

- Acharya, S. and S. S. Karim (1996). "Modulation of GDP release from transducin by the conserved Glu134-Arg135 sequence in rhodopsin." *J Biol Chem* 271(41): 25406-11.
- Alvewjse, A. E., H. Timmerman, et al. (2000). "The Effect of Mutations in the DRY Motif on the Constitutive Activity and Structural Instability of the Histamine H(2) Receptor." *Mol Pharmacol* 57(3): 890-898.
- Allen, L. F., R. J. Lefkowitz, et al. (1991). "G-protein-coupled receptor genes as protooncogenes: constitutively activating mutation of the alpha 1B-adrenergic receptor enhances mitogenesis and tumorigenicity." *Proc Natl Acad Sci USA* 88(24): 11354-8.
- Amaludru, T. T., Jrd, S. Drages-Gronio, et al. (1995). "Signal transduction by the formyl peptide receptor. Studies using chimeric receptors and site-directed mutagenesis define a novel domain for interaction with G-proteins." *J Biol Chem* 270(47): 28010-3.
- Bhimi, K., E. Mutschler, et al. (1994). "Functional role in ligand binding and receptor activation of an asparagine residue present in the sixth transmembrane domain of all muscarinic acetylcholine receptors." *J Biol Chem* 269(29): 18870-6.
- Boone, C. N., G. Davis, et al. (1993). "Mutations that alter the third cytoplasmic loop of the α -factor receptor lead to a constitutive and hypersensitive phenotype." *Proc Natl Acad Sci USA* 90(21): 9921-5.
- Bradbury, F. A., N. Kawate, et al. (1997). "Post-translational processing in the Golgi plays a critical role in the trafficking of the luteinizing hormone/human chorionic gonadotropin receptor to the cell surface." *J Biol Chem* 272(9): 5921-6.
- Bradbury, F. A. and K. M. Menon (1999). "Evidence that constitutively active luteinizing hormone/human chorionic gonadotropin receptors are rapidly internalized." *Biochemistry* 38(27): 8703-12.
- Burger, M., J. A. Burger, et al. (1999). "Point mutation causing constitutive signaling of chimeric m2/m5 muscarinic receptors and delineation of G-protein coupling selectivity domains." *coupled receptor." J Immunol* 163(4): 2017-22.
- Burstein, E. S., T. A. Spalding, et al. (1996). "Constitutive activation of chimeric m2/m5 muscarinic receptors and delineation of G-protein coupling selectivity domains." *Biochem Pharmacol* 51(4): 539-44.
- Caavalli, A., A. M. Babey, et al. (1999). "Altered adenylyl cyclase responsiveness subsequent to point mutations of Asp 128 in the third transmembrane domain of the delta-opioid receptor." *Neuroscience* 93(3): 1025-31.
- Chapronier, S., K. R. Jarvie, et al. (1996). "Silencing of the constitutive activity of the dopamine D1B receptor. Reciprocal mutations between D1 receptor subtypes delineate residues underlying activation properties." *J Biol Chem* 271(45): 28071-6.
- Cho, W., L. P. Taylor, et al. (1996). "Mutagenesis of residues adjacent to transmembrane prolines alters D1 dopamine receptor binding and signal transduction." *Mol Pharmacol* 50(5): 1338-45.
- Cohen, D. P., C. N. Thaw, et al. (1997). "Human calcitonin receptors exhibit agonist-independent (constitutive) signaling activity." *Endocrinology* 138(4): 1400-5.
- Cohen, G. B., T. Yang, et al. (1993). "Constitutive activation of opsin: influence of charge at position 134 and size at position 296." *Biochemistry* 32(23): 6111-5.
- Dube, P., A. DeCostanzo, et al. (2000). "Interaction between transmembrane domains five and six of the alpha 1A-factor receptor." *J Biol Chem* 275(34): 26492-9.
- Duprez, L., J. Parma, et al. (1994). "Germline mutations in the thyrotropin receptor gene cause non-autoimmune autoimmune dominant hyperthyroidism." *Nat Genet* 7(3): 396-401.
- Egan, C. T., K. Herrick-Davis, et al. (1998). "Creation of a constitutively activated state of the 5-hydroxytryptamine2A receptor by site-directed mutagenesis: inverse agonist activity of antipsychotic drugs." *J Pharmacol Exp Ther* 284(1): 83-90.
- Esapa, C. T., L. Duprez, et al. (1999). "A novel thyrotropin receptor mutation in an infant with severe thyrotoxicosis." *Thyroid* 9(10): 1005-10.
- Fanelli, F., P. Barbier, et al. (1999). "Activation mechanism of human oxytocin receptor: a combined study of experimental and computer-simulated mutagenesis." *Mol Pharmacol* 56(1): 214-25.
- Gaudin, P., J. J. Maoret, et al. (1998). "Constitutive activation of the human vasopressin receptor 1 receptor, a member of the new class II family of G protein-coupled receptors." *J Biol Chem* 273(9): 4990-6.
- Gaudin, P., C. Rouyer-Fessard, et al. (1998). "Constitutive activation of the human VIP1 receptor." *Ann NY Acad Sci* 865: 382-5.

FIG. 1 (14 of 15)

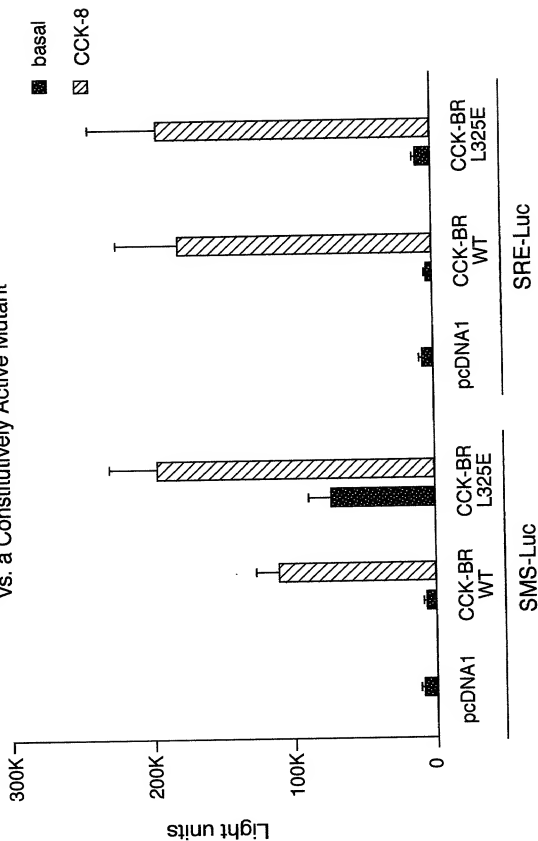
- Gowarthan, C. P. and D. D. Oprea (1994). "Active site-directed inactivation of constitutively active mutants of rhodopsin." *J Biol Chem* 269(9): 6524-7.
- Grobleski, T. B., Maigret, et al. (1997). "Mutation of Asn111 in the third transmembrane domain of the AT1A angiotensin II receptor induces its constitutive activation." *J Biol Chem* 272(3): 1822-6.
- Han, M., S. O. Smith, et al. (1998). "Constitutive activation of opsin by mutation of methionine 257 on transmembrane helix 6." *Biochemistry* 37(22): 8253-61.
- Haegawa, H., M. Negishi, et al. (1996). "Two isoforms of the prostaglandin E receptor EP3 subtype differ in agonist-independent constitutive activity." *J Biol Chem* 271(4): 1857-60.
- Herrick-Davis, K., C. Egan, et al. (1997). "Activating mutations of the serotonin 5-HT2C receptor." *J Neurochem* 69(3): 1138-44.
- Fjorft, S. A., C. Orskov, et al. (1998). "Constitutive activity of glucagon receptor mutants." *Mol Endocrinol* 12(1): 78-86.
- Hogger, P., M. S. Shockey, et al. (1995). "Activating and inactivating mutations in N- and C-terminal B loop junctions of muscarinic acetylcholine Hm1 receptors." *J Biol Chem* 270(13): 7405-10.
- Hwa, J., R. Izumi, et al. (1997). "Synergism of constitutive activity in alpha 1-adrenergic receptor activation." *Biochemistry* 36(3): 633-9.
- Ishii, I., T. Ganai, et al. (1997). "Alanine exchanges of polar amino acids in the transmembrane domains of a platelet-activating factor receptor generate both constitutively active and inactive mutants." *J Biol Chem* 272(12): 7846-54.
- Jensen, A. A., T. A. Spalding, et al. (2000). "Functional importance of the Ala116-Pro136 region in the calcium-sensing receptor. CONSTITUTIVE ACTIVITY AND INVERSE ALGONISM IN A FAMILY C-G-PROTEIN-COUPLED RECEPTOR [In Process Citation]." *J Biol Chem* 275(38): 29347-55.
- Jin, J., G. F. Mao, et al. (1997). "Constitutive activity of human prostaglandin E receptor EP3 isoforms." *British J Pharmacol* 121: 317-23.
- Kjalsberg, M. A., S. Cotecchia, et al. (1992). "Constitutive activation of the alpha 1B-adrenergic receptor by all amino acid substitutions at a single site. Evidence for a region which constrains receptor activation." *J Biol Chem* 267(3): 1430-3.
- Konopka, J. B., S. M. Margalit, et al. (1996). "Mutation of Pro-258 in transmembrane domain 6 constitutively activates the G protein-coupled alpha-factor receptor." *Proc Natl Acad Sci USA* 93(13): 6764-9.
- Kouyama, S., C. Van Dop, et al. (1995). "Characterization of heterogeneous mutations causing constitutive activation of the human luteinizing hormone receptor in familial male precocious puberty." *Hum Mol Genet* 4(2): 183-8.
- Kudo, M., Y. Otsuga, et al. (1996). "Transmembrane regions V and VI of the human luteinizing hormone receptor are required for constitutive activation by a mutation in the third intracellular loop." *J Biol Chem* 271(37): 22470-8.
- Lefkowitz, R. J., S. Cotecchia, et al. (1993). "Constitutive activity of receptors coupled to guanine nucleotide regulatory proteins." *Trends Pharmacol Sci* 14(8): 303-7.
- Lin, J., N. Blin, et al. (1996). "Molecular mechanisms involved in muscarinic acetylcholine receptor-mediated G protein activation studied by insertion mutagenesis." *J Biol Chem* 271(11): 6172-8.
- Marin, J., C. Koch, et al. (1999). "Constitutive activation of the human bradykinin B2 receptor induced by mutations in transmembrane helices III and VI." *Mol Pharmacol* 55(1): 92-101.
- Mason, D. A., J. D. Moore, et al. (1999). "A gain-of-function polymorphism in a G-protein coupling domain of the human beta1-adrenergic receptor." *J Biol Chem* 274(18): 12670-4.
- Maus-Luehovich, N., D. R. Nussenzweig, et al. (1995). "Truncation of the thyrotropin-releasing hormone receptor carboxyl tail causes constitutive activity and leads to impaired responsiveness in Xenopus oocytes and A120 cells." *J Biol Chem* 270(3): 1041-7.
- Morin, D., N. Condeelis, et al. (1998). "The D136A mutation of the V2 vasopressin receptor induces a constitutive activity which permits discrimination between antagonists with partial agonist and inverse agonist activities." *FEBS Lett* 441(3): 470-5.
- Narverez, T., L. Wang, et al. (1996). "Thrombin receptor activating mutations. Alteration of an extracellular agonist recognition domain causes constitutive signaling." *J Biol Chem* 271(2): 702-6.
- Olesnicki, N. S., A. J. Brown, et al. (1999). "A constitutively active G-protein-coupled receptor causes mating self-compatibility in the mushroom *Coprinus*." *Embo J* 18(10): 2756-63.
- Parent, J. L., C. Le Guillou, et al. (1996). "Mutations of two adjacent amino acids generate inactive and constitutively active forms of the human platelet-activating factor receptor." *J Biol Chem* 271(14): 7949-55.

FIG. 1 (15 of 15)

- Parma, J., J. Van Sande, et al. (1995). "Somatic mutations causing constitutive activity of the thyrotropin receptor are the major cause of hyperfunctioning thyroid adenomas: identification of additional mutations activating both the cyclic adenosine 3',5'-monophosphate and inositol phosphate-Ca²⁺ cascades." *Mol. Endocrinol.* 9(6): 725-33.
- Parnot, C., S. Bardin, et al. (2000). "Systematic identification of mutations that constitutively activate the thyrotropin receptor by screening a randomly mutated cDNA library with an original pharmacological bioassay." *Proc Natl Acad Sci USA* 97(13): 7615-20.
- Paschke, R., M. Tonacchera, et al. (1994). "Identification and functional characterization of two new somatic mutations causing constitutive activation of the thyrotropin receptor in hyperfunctioning autonomous adenomas of the thyroid." *J Clin Endocrinol Metab* 79(6): 1785-9.
- Pawels, P., J. A. Gouble, et al. (1999). "Activation of constitutive 4-hydroxyphenylamine B receptor by a series of mutations in the BBXXB motif: positioning of the third intracellular loop distal junction and its goipha protein interactions [in Process Citation]." *Biochem J* 343 Pt 2: 435-42.
- Perez, D. M., J. Hwa, et al. (1996). "Constitutive activation of a single effector pathway: evidence for multiple activation states of a G protein-coupled receptor." *Mol Pharmacol* 49(1): 112-22.
- Ren, Q., H. Kurose, et al. (1993). "Constitutively active mutants of the alpha 2-adrenergic receptor [published erratum appears in J Biol Chem 1994 Jan 14;269(2):1566]." *J Biol Chem* 268(22): 16483-7.
- Rim, J. and D. D. Orian (1993). "Constitutive activation of opsin: interaction of mutants with rhodopsin kinase and arrestin." *Biochemistry* 34(37): 11938-45.
- Robbins, L. S., J. H. Nadeau, et al. (1993). "Pigmentation phenotypes of variant extension locus alleles result from point mutations that alter MSH receptor function." *Cell* 72(6): 827-34.
- Russo, D., M. G. Wong, et al. (1999). "A Val 677 activating mutation of the thyrotropin receptor in a Hurthle cell thyroid carcinoma associated with thyrotoxicosis." *Thyroid* 9(1): 13-7.
- Samama, P., S. Cotecchia, et al. (1993). "A mutation-induced activated state of the beta 2-adrenergic receptor: Extending the ternary complex model." *Journal of Biological Chemistry* 268(7): 4625-36.
- Scheer, A., T. Costa, et al. (2000). "Mutational analysis of the highly conserved arginine within the Glu/Asp-Arg-Tyr motif of the alpha (b)-adrenergic receptor: effects on receptor isomerization and activation." *Mol Pharmacol* 57(2): 219-31.
- Scheer, A., F. Fanelli, et al. (1997). "The activation process of the alpha 1B-adrenergic receptor: potential role of protonation and hydrophobicity of a highly conserved aspartate." *Proc Natl Acad Sci USA* 94(3): 808-13.
- Schipani, E., G. S. Jensen, et al. (1997). "Constitutive activation of the cyclic adenosine 3',5'-monophosphate signaling pathway by parathyroid hormone (PTH)/PTH-related peptide receptors mutated at the two loci for Jansen's metaphyseal chondrodysplasia." *Mol Endocrinol* 11(7): 851-8.
- Shenker, A., L. Laue, et al. (1993). "A constitutively activating mutation of the luteinizing hormone receptor in familial male precocious puberty [see comments]." *Nature* 365(6447): 652-4.
- Sommers, C. M., N. P. Martin, et al. (2000). "A limited spectrum of mutations causes constitutive activation of the yeast alpha-factor receptor." *Biochemistry* 39(23): 6898-909.
- Spalding, T. A., E. S. Brustein, et al. (1998). "Identification of a ligand-dependent switch within a muscarinic receptor." *J Biol Chem* 273(34): 21563-8.
- Spalding, T. A., E. S. Brustein, et al. (1997). "Constitutive activation of the m5 muscarinic receptor by a series of mutations at the extracellular end of transmembrane 6." *Biochemistry* 36(33): 10109-16.
- Tsang, C. C. and L. Lin (1997). "A point mutation in the glucose-dependent insulinotropic peptide receptor confers constitutive activity." *Biochem Biophys Res Commun* 232(1): 96-100.
- Wonerow, P., T. Schoneberg, et al. (1998). "Deletions in the third intracellular loop of the thyrotropin receptor. A new mechanism for constitutive activation." *J Biol Chem* 273(14): 7900-5.

FIG. 2

Light Emission Induced by the WT CCK-BR
vs. a Constitutively Active Mutant



17/27

FIG. 3

A Point Mutation Confers Constitutive Activity
to the Rat μ Opioid Receptor

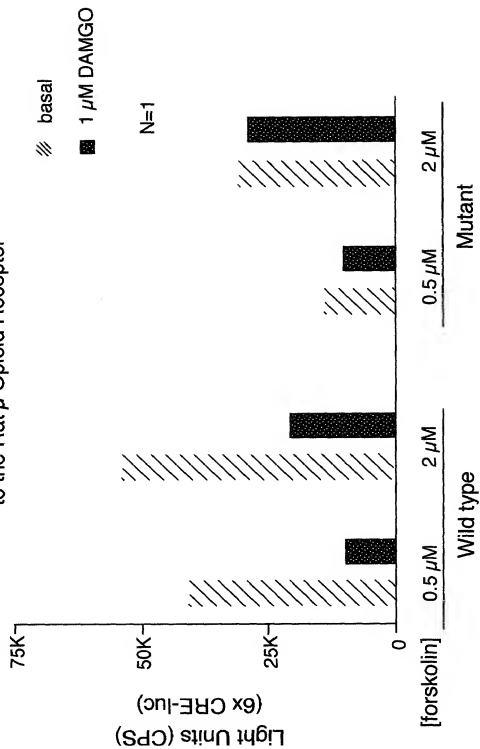


FIG. 4

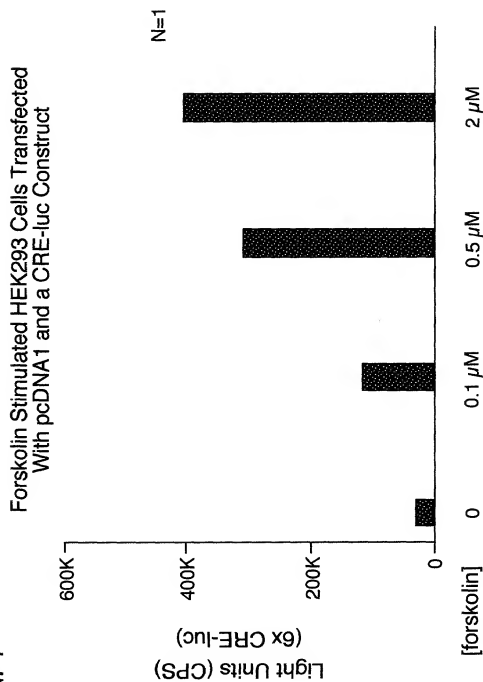
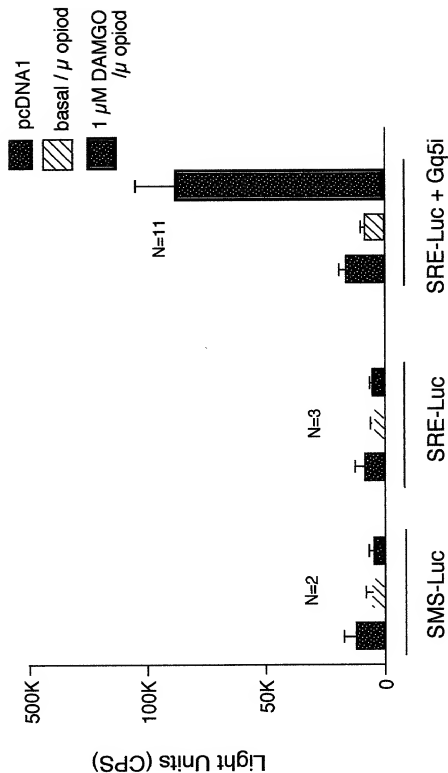


FIG. 5

The Rat μ Opioid Receptor Signals Through $G\alpha i$ 

20/27

FIG. 6

A Point Mutation Confers Constitutive Activity to the Rat μ Opioid Receptor

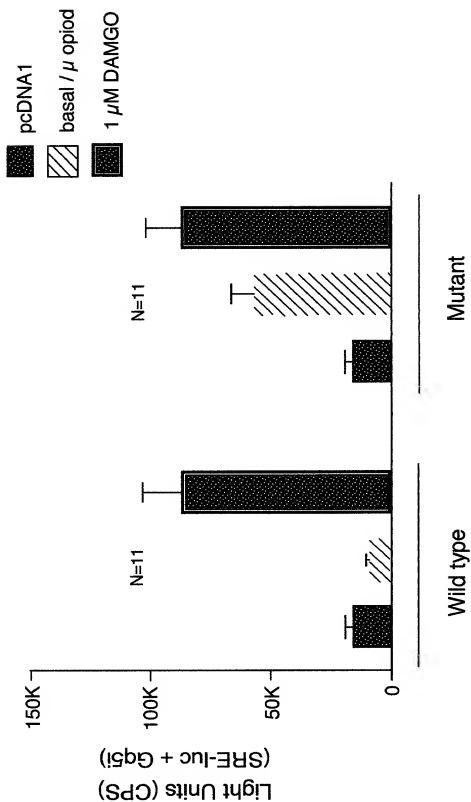
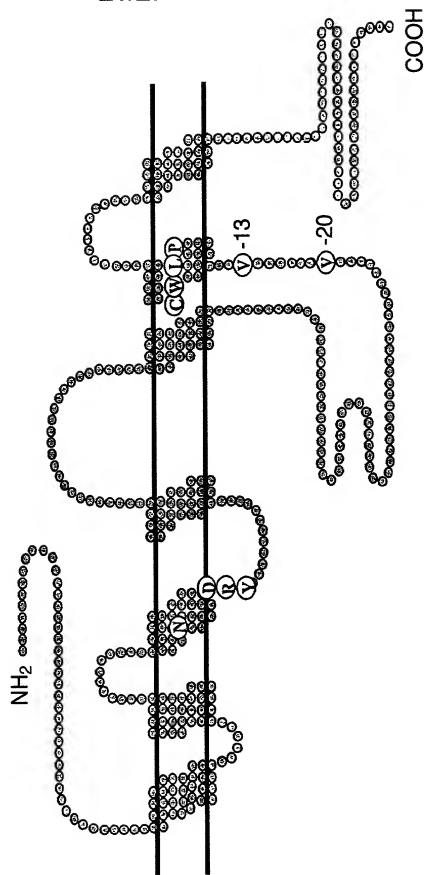


FIG. 7

Target Residues Within Class I GPCR's



22/27

FIG. 8

TMD III Asn (-14 from DRY) is a Target
for Mutation Induced Constitutive Activity

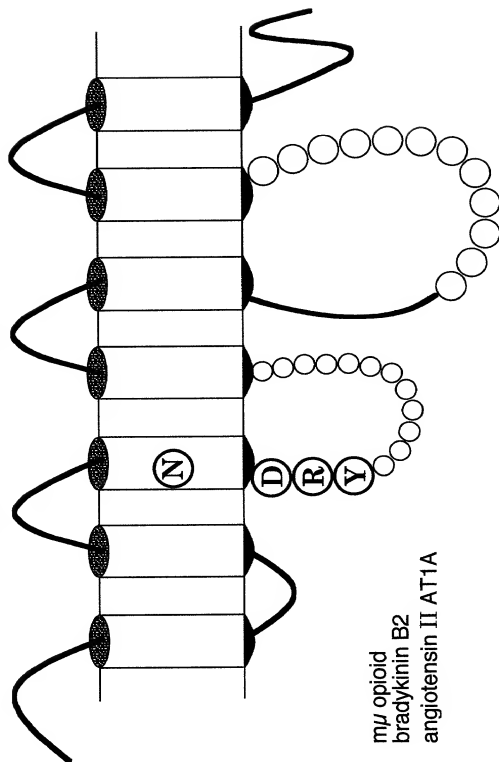
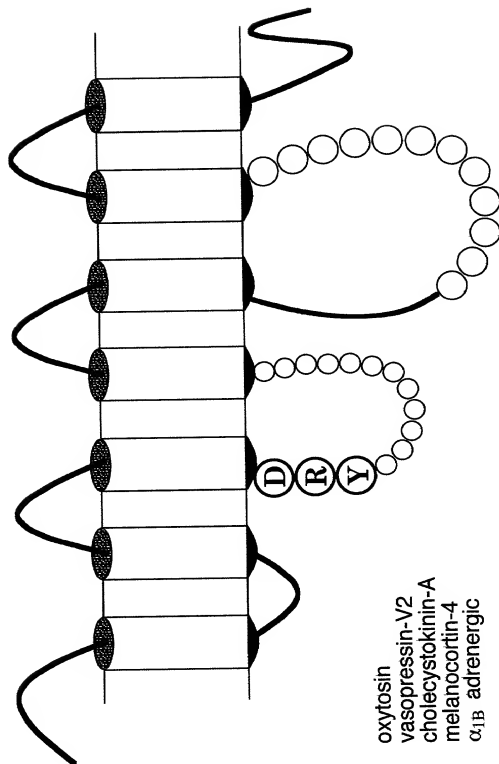


FIG. 9

The 'DRY' Motif is a Target for Mutation
Induced Constitutive Activity



24/27

FIG. 10

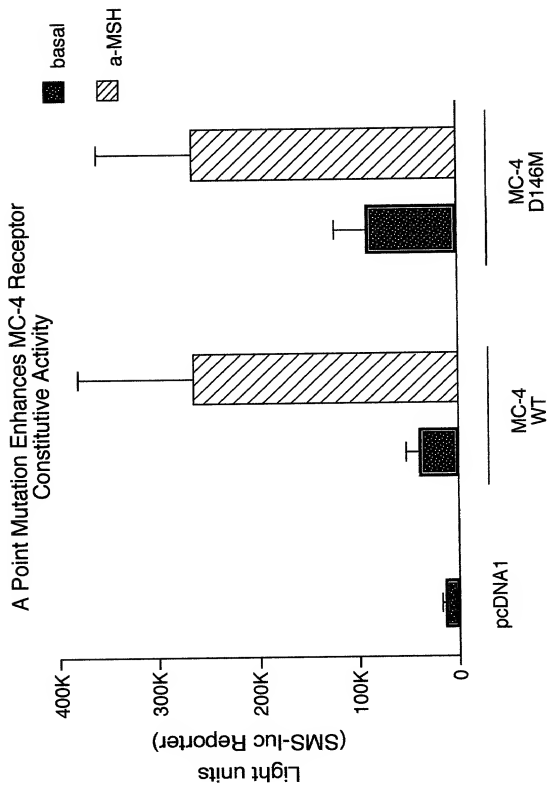


FIG. 11

The -13 Position is a Target for Mutation
Induced Constitutive Activity

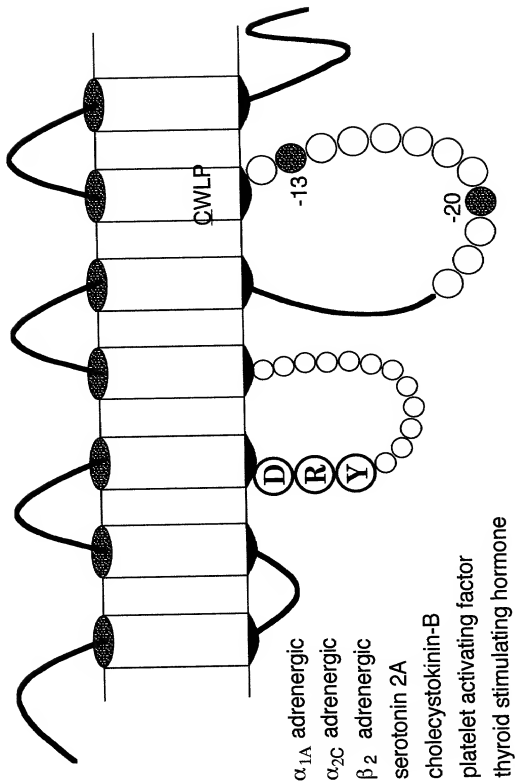


FIG. 12

26/27

```

ork 1 -----MESI FRGEPGETCAISACTPPNSESAMFPGWAEF INGSAGSDEDAE
orkr 1 -----MESPIQIFRGEPEGETCAISACTLPNSESAMFPGWAESS..DSNGSVGSEDDQ
orm 1 MDSSAAPTINASNTDABAYSSCSAPSPSGSW..NLSHLDNLSPGCGPNRTDLGGNDEL
ormr 1 MDSSGTGPGNTSDCSDPFAQASCSPA..PGSW..NLSHVDENOSPCCGLNRTGLGNDL
ord 1 -----MEPAPSAGAE..PPLFPMASAPSPACSPASANASG
ATla 1 -----MALNSSAEDGIKRIS
BK-2 1 -----MFSPWKISMFLSVREDSVPTTAFSRLMLNVTLLQGPTLNG..TFAQ

```

```

ork 49 LEPAHISEAH..PVAITAHYSVVFVUGLNGSVAHVIITRYTKMTATNIYIFNLALADA
orkr 49 LEPAHISEAH..PVAITAHYSVVFVUGLNGSVAHVIITRYTKMTATNIYIFNLALADA
orm 59 CPPTGS..ESMTAHTIMALYSHVCVVGLFGMFLVAVHVIITRYTKMTATNIYIFNLALADA
ormr 57 CPQTGS..ESMTAHTIMALYSHVCVVGLFGMFLVAVHVIITRYTKMTATNIYIFNLALADA
ord 37 PPGASASSIALARATATYASVAVCVGLFGMFLVAVHVIITRYTKMTATNIYIFNLALADA
ATla 16 DDCPAGRHSYIFVHPTLYSHVGLFGMFLVAVHVIITRYTKMTATNIYIFNLALADA
BK-2 45 SKCPQVWELGLWNTLQPPFLWVHVFVATBEHIFVLSVFLCHKSSCVAEITVGLHFLAADI

```

```

ork 107 IVTHITPEFOSVYVLNN..SWPFGHLCRIVISIDYYNMETSIFLITLMSVDRYIAVCHPVV
orkr 107 IVTHITPEFOSAVYVLNN..SWPFGHLCRIVISIDYYNMETSIFLITLMSVDRYIAVCHPVV
orm 118 LAISTITPEFOSAVYVLNG..SWPFGHLCRIVISIDYYNMETSIFLITLMSVDRYIAVCHPVV
ormr 116 LAISTITPEFOSAVYVLNG..SWPFGHLCRIVISIDYYNMETSIFLITLMSVDRYIAVCHPVV
ord 97 LAISTITPEFOSAKYLAKE..SWPFGHLCRIVISIDYYNMETSIFLITLMSVDRYIAVCHPVV
ATla 76 CFLITLPELVYTAIEYKMEPEGNLCIASSSVTEHLSAFLHLCSSADRYIAVCHPVV
BK-2 105 ILACGDFEWATFISNNFELSGEITLCRAVNHISMSSTICLMAUSDRYIAVCHPVV

```

-14 from DRY

```

ork 166 ALDERTELKAKIINICINILASSVCHSAIVLGGTKVR..EDVDITCEHLOPDDDSYAMD
orkr 166 ALDERTELKAKIINICINILASSVCHSAIVLGGTKVR..EDVDITCEHLOPDDDSYAMD
orm 177 ALDERTELKAKIINICINILASSVCHSAIVLGGTKVR..EDVDITCEHLOPDDDSYAMD
ormr 175 ALDERTELKAKIINICINILASSVCHSAIVLGGTKVR..EDVDITCEHLOPDDDSYAMD
ord 156 ALDERTELKAKIINICINILASSVCHSAIVLGGTKVR..EDVDITCEHLOPDDDSYAMD
ATla 136 SRLRLMLVAKYTCIITLWAGLASDEAVIHRNV..YFINTNTVCAFHYESRN..STLP
BK-2 165 MGRMEGVRWAKYYSVIVIGCLLSSSENVFRTMKEYSDCHNVYACVHESMS...LIME

```

```

ork 224 EFMICVFIAPVLPVLLITVGYTLMILRLKSVRLSGSHEKDRNLRLRITRVLVVVAVE
orkr 224 EFMICVFIAPVLPVLLITVGYTLMILRLKSVRLSGSHEKDRNLRLRITRVLVVVAVE
orm 232 NLMICVFIAPVLPVLLITVGYTLMILRLKSVRLSGSHEKDRNLRLRITRVLVVVAVE
ormr 230 NLMICVFIAPVLPVLLITVGYTLMILRLKSVRLSGSHEKDRNLRLRITRVLVVVAVE
ord 211 TVTICVFIAPVLPVLLITVGYTLMILRLKSVRLSGSHEKDRNLRLRITRVLVVVAVE
ATla 193 EFMICVFIAPVLPVLLITVGYTLMILRLKSVRLSGSHEKDRNLRLRITRVLVVVAVE
BK-2 222 VETNMLNVVGFELP..LSVITFCINCHQLNRNNEKQKRIQTE..RRHVLVVLVVLLE

```

```

ork 284 IVCWTPDHITFVVAELGS..T....SHETALSSNYECIALGYTNSCNPLVYAFIDENF
orkr 284 IVCWTPDHITFVVAELGS..T....SHETALSSNYECIALGYTNSCNPLVYAFIDENF
orm 292 IVCWTPDHITFVVAELGS..T....SHETALSSNYECIALGYTNSCNPLVYAFIDENF
ormr 290 IVCWTPDHITFVVAELGS..T....SHETALSSNYECIALGYTNSCNPLVYAFIDENF
ord 271 VVCWTPDHITFVVAELGS..T....SHETALSSNYECIALGYTNSCNPLVYAFIDENF
ATla 250 FFSVWGHOTFTFVWVGHGVHIDCKI..SDIDTMTPTICTVYFVNCINPLVYAFIDENF
BK-2 280 IVCWTPDHITFVVAELGS..T....SHETALSSNYECIALGYTNSCNPLVYAFIDENF

```

SEQ ID NO:

```

ork 338 KRCFREFCFPIKMREROSTSRNR..NTYOD..PAYLRDMDGYNKPY----- 76
orkr 338 KRCFREFCFPIKMREROSTSRNR..NTYOD..PAMEDVGGYKPY----- 77
orm 346 KRCFREFCFPISSNHCQSTSRNR..NTYOD..PAMEDVGGYKPY----- 78
ormr 344 KRCFREFCFPISSNHCQSTSRNR..NTYOD..PAMEDVGGYKPY----- 79
ord 326 KRCFREFCFPISSNHCQSTSRNR..NTYOD..PAMEDVGGYKPY----- 80
ATla 310 KRYELQLLKYPKPKAKSHS...SLSTKM..STLEYRPSNMSSSAKKPASCPEVE- 81
BK-2 340 KRKSWVYVGVCGGGRSEPIQMESEK..GTL..RTSISVPECHKIQWAGRSQ 82

```

00966871.031202

mORmouse 1 MDSSAGGGMISDCSDPDA..PASCSPA..EGSWHMLSHRDGMSDPOGNNRVLGCGSHSLG
 mORrat 1 MDSSITGGGMSDSCSDPDA..QASCSPA..EGSWHMLSHRDGMSDPOGNNRVLGCGSHSLG
 mORbovin 1 MDSSGVPTWASGNDIDFTHPSSCSPPAPSSSSAMFSSDAGMLSDPOGNNRVLGCGSHSLG
 mORhuman 1 MDSSAGPTWASGNDIDAY..SCSPAPSSSGSAMFSSDAGMLSDPOGNNRVLGCGSHSLG
 mORpig 1 MDSSAGPTWASGNDIDFSPSSMSSEVPSSSAMFSSDAGMLSDPOGNNRVLGCGSHSLG
 mORws 1 MESS..GMSDFLYPLS.....MVMG....NSSVLGRFSSSTSTFLNMNGSSSDTD
 Atla 1MALNSAETCHRIQDDC
 BK-2 1 -----MFSPNKISMFLVREDSVPTTASFADMLNVLTGPTLNG..TFACSK

mORmouse 58 PONGSPSNATLITKALYSIVCVGLGCMELVWVIVRYTKMKPATNTIENIALADALA
 mORrat 58 PONGSPSNATLITKALYSIVCVGLGCMELVWVIVRYTKMKPATNTIENIALADALA
 mORbovin 61 PONGSPSNATLITKALYSIVCVGLGCMELVWVIVRYTKMKPATNTIENIALADALA
 mORhuman 60 PONGSPSNATLITKALYSIVCVGLGCMELVWVIVRYTKMKPATNTIENIALADALA
 mORpig 61 PONGSPSNATLITKALYSIVCVGLGCMELVWVIVRYTKMKPATNTIENIALADALA
 mORws 61 POKKS..TITLITTTATGNSVGLV..TAAATATREVTATNTIENIALADALA
 Atla 1 EKASRHSITPVE..PTLISHPVCHRETSVAILIYFYFKKVASNRILALALCLF
 BK-2 48 EOVENLGWNTII..QPFLMVLVPTLEMLFVPSFCLIKSSCVLAETGCGIDAPAILI

mORmouse 118 TSTAPROSVNYLAG..TWBPCNIIICQVVISIDYNNFSTIETLCTMSVDRYIAVCHPVIAL
 mORrat 118 TSTAPROSVNYLAG..TWBPCNIIICQVVISIDYNNFSTIETLCTMSVDRYIAVCHPVIAL
 mORbovin 121 TSTAPROSVNYLAG..TWBPCNIIICQVVISIDYNNFSTIETLCTMSVDRYIAVCHPVIAL
 mORhuman 120 TSTAPROSVNYLAG..TWBPCNIIICQVVISIDYNNFSTIETLCTMSVDRYIAVCHPVIAL
 mORpig 121 TSTAPROSVNYLAG..TWBPCNIIICQVVISIDYNNFSTIETLCTMSVDRYIAVCHPVIAL
 mORws 121 TSTAPROSVNYLAG..TWBPCNIIICQVVISIDYNNFSTIETLCTMSVDRYIAVCHPVIAL
 Atla 78 LLNLEWNYTAKYRREPCNIIICQVVISIDYNNFSTIETLCTMSVDRYIAVCHPVIAL
 BK-2 107 ACCGPMATITISNFDLPGETLCPVWITLISYSSHCFPMALGSDRYIAVCHPVIAL

mORmouse 177 DERTPRNAKINWCVNITLSSAIGLPMVMATTKYRC.....GSIDCLTFSHPTWYME
 mORrat 177 DERTPRNAKINWCVNITLSSAIGLPMVMATTKYRC.....GSIDCLTFSHPTWYME
 mORbovin 180 DERTPRNAKINWCVNITLSSAIGLPMVMATTKYRC.....GSIDCLTFSHPTWYME
 mORhuman 179 DERTPRNAKINWCVNITLSSAIGLPMVMATTKYRC.....GSIDCLTFSHPTWYME
 mORpig 180 DERTPRNAKINWCVNITLSSAIGLPMVMATTKYRC.....GSIDCLTFSHPTWYME
 mORws 166 DERTPRNAKINWCVNITLSSAIGLPMVMATTKYRC.....GSIDCLTFSHPTWYME
 Atla 138 LKRLMLVATCHIIEAGLASIEPWHRV...YFIENTNITVCAPHRESNSTLP
 BK-2 167 RMGVNEMATKYSLEVIIGCHLLSSHPVFRME.....EYSDEGHNVTAQVSTPS...LINE

mORmouse 230 NLLIKCVFIAPIMEVLITTCVGLMILRLKSVNLSGSKEDRNLRLRITRMVLVVAVF
 mORrat 230 NLLIKCVFIAPIMEVLITTCVGLMILRLKSVNLSGSKEDRNLRLRITRMVLVVAVF
 mORbovin 233 NLLIKCVFIAPIMEVLITTCVGLMILRLKSVNLSGSKEDRNLRLRITRMVLVVAVF
 mORhuman 232 NLLIKCVFIAPIMEVLITTCVGLMILRLKSVNLSGSKEDRNLRLRITRMVLVVAVF
 mORpig 233 NLLIKCVFIAPIMEVLITTCVGLMILRLKSVNLSGSKEDRNLRLRITRMVLVVAVF
 mORws 226 TLAIKCVFIAPIMEVLITTCVGLMILRLKSVNLSGSKEDRNLRLRITRMVLVVAVF
 Atla 193 IGGGTINLGEITFATLTSITLTKALSKAYEHONNPRNDS...FRATLALFLF
 BK-2 222 VFTNMLNIVGDEP..LSHITFCTCKHOLLRNNEVOKPTEIQTE..RRAVLVVLVVL

mORmouse 290 IVCWTPHIVVILKALTI.....PETTEQTVSWHEFCIALGYTNSCLNPVLYAFIDENF
 mORrat 290 IVCWTPHIVVILKALTI.....PETTEQTVSWHEFCIALGYTNSCLNPVLYAFIDENF
 mORbovin 293 IVCWTPHIVVILKALTI.....PETTEQTVSWHEFCIALGYTNSCLNPVLYAFIDENF
 mORhuman 292 IVCWTPHIVVILKALTI.....PETTEQTVSWHEFCIALGYTNSCLNPVLYAFIDENF
 mORpig 293 IVCWTPHIVVILKALTI.....PETTEQTVSWHEFCIALGYTNSCLNPVLYAFIDENF
 mORws 286 IVCWTPHIVVILKALTI.....ENSLQVWHEFCIALGYTNSCLNPVLYAFIDENF
 Atla 250 FFSVWHEFCIALGYTNSCLNPVLYAFIDENF
 BK-2 280 HVCWTPHIVVILKALTI.....ENSLQVWHEFCIALGYTNSCLNPVLYAFIDENF

SEQ ID NO:

mORmouse 344 KRCSREFC..LPTSSITDQNSARIRONTREHPSTANTVDTRNHQLENLEAEATAPLE 83
 mORrat 344 KRCSREFC..LPTSSITDQNSARIRONTREHPSTANTVDTRNHQLENLEAEATAPLE 79
 mORbovin 347 KRCSREFC..LPTSSITDQNSARIRONTREHPSTANTVDTRNHQLENLEAEATAPLE 84
 mORhuman 346 KRCSREFC..LPTSSITDQNSARIRONTREHPSTANTVDTRNHQLENLEAEATAPLE 85
 mORpig 347 KRCSREFC..LPTSSITDQNSARIRONTREHPSTANTVDTRNHQLENLEAEATAPLE 86
 mORws 340 KRCSREFC..LPTSSITDQNSARIRONTREHPSTANTVDTRNHQLENLEAEATAPLE 87
 Atla 310 KRYVLLKLYIKPKKHS..SLSTKSTLYRPSDNGSSAKKPCASCFEVE----- 81
 BK-2 340 RKSWSVYQGCCKGGRCSPIQMENSMGLT..RPSISVEROIEKQDWAESQRO--- 82